



SITE LOGIC Report

Stable Isotope Probing (SIP)

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Report Date: July 8, 2020

Project: FWAFB ST012, 9101110001.5310.02

Comments:

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Executive Summary

A Stable Isotope Probing (SIP) study was performed to determine whether biodegradation of benzene is occurring under existing site conditions. Bio-Trap® samplers baited with ¹³C-labeled benzene were deployed in UWBZ26 033020, UWBZ27 033020, LSBZ38 033020 and LSBZ39 033020. Following a 35-day deployment period, the Bio-Traps were recovered to quantify ¹³C incorporation into biomass and dissolved inorganic carbon (DIC). A complete summary of the SIP results is provided in Table 1 and Figures 1 through 5. Following are the key observations from the results obtained for the monitoring wells.

Stable Isotope Probing Results

- Quantification of 13 C-enriched PLFA conclusively demonstrated that benzene was metabolized under existing site conditions. The average PLFA δ^{13} C value in all four wells fell within the moderate range, indicating 13 C-labeled benzene was incorporated into microbial biomass.
- The DIC δ^{13} C value in UWBZ26 033020 measured at 38‰, although low, this indicates that a small amount of benzene was also mineralized during the deployment period. The DIC values for UWBZ27 033020, LSBZ38 033020 and LSBZ39 033020 were near background levels, indicating that the mineralization of benzene is limited under current site conditions for these wells.
- The total PLFA biomass concentration in UWBZ26 03302 was on the order of 10⁷ cells/bead, which was within the high range while the biomass concentrations for UWBZ27 033020, LSBZ38 033020 and LSBZ39 033020 fell within the moderate range at 10⁵ cells/bead.
- The PLFA community structures for all samples were primarily composed of indicators of proteobacteria and normal saturates with UWBZ26 033020 also having a high percentage of eukaryote indicators. Indicators of firmicutes were present in samples UWBZ27 033020, LSBZ38 033020 and LSBZ39 033020 and to a lower extent in UWBZ26 033020. Indicators of actinomycetes and anaerobic metal reducers were only present in UWBZ27 033020. Eukaryote indicators were also present in this sample.



Overview of Approach

Stable Isotope Probing (SIP)

Stable isotope probing (SIP) is an innovative approach to conclusively determine whether *in situ* biodegradation of a contaminant of concern is occurring.

With the SIP method, a Bio-Trap® is amended with a specially synthesized ¹³C form of the contaminant of concern (e.g. ¹³C-benzene). The ¹³C essentially serves as a "label" to track biodegradation. For petroleum hydrocarbons and many other contaminants, biodegradation is a process whereby some microorganisms use the contaminant of concern as a carbon and energy source. When used as carbon source, contaminant carbon is incorporated into biomolecules such as phospholipids, DNA, and proteins supporting growth of new cells (biomass). When used as an energy source, contaminant carbon is oxidized to CO₂ as part of cellular metabolism. Thus, detection of the ¹³C "label" in the end products of biodegradation (bacterial biomass and CO₂) at the end of the SIP study provides conclusive evidence of contaminant biodegradation.

To perform a SIP study, a Bio-Trap® is amended with the ¹³C form of the contaminant of concern (e.g. ¹³C-benzene) and deployed in an existing monitoring well for a period of 30 to 60 days. If present and active under the existing subsurface conditions, bacteria capable of utilizing the ¹³C labeled contaminant of concern will colonize and grow in the Bio-Trap® over the course of the deployment period. Following recovery from the well, the Bio-Trap® is shipped to the laboratory and two approaches are used to conclusively evaluate contaminant biodegradation:

- Quantification of ¹³C enriched phospholipid fatty acids (PLFA)
- Quantification of ¹³C enriched dissolved inorganic carbon (DIC)

PLFA are a primary component of the membrane of bacterial cells and have long been used as a measure of microbial biomass. The detection of ¹³C enriched PLFA during a SIP study indicates incorporation into microbial biomass and therefore conclusively demonstrates contaminant biodegradation.

Detection of ¹³C enriched DIC which includes ¹³CO₂ conclusively indicates contaminant biodegradation and mineralization.



Results

Table 1. Summary of the stable isotope probing results obtained from the Bio-Trap® Units.

Sample Name	UWBZ26 033020	UWBZ27 033020	LSBZ38 033020	LSBZ39 033020
Sample Date	3/30/2020	3/30/2020	3/30/2020	3/30/2020
MHD	122RC1	122RC2	122RC3	1922R@4
¹³ C Contaminant Loss				
13C Benzene Pre-deployment (µg/bead)	177 ± 3	177 ± 3	177 ± 3	177 ± 3
¹³ C Benzene Post-deployment (μg/bead)	2 ± 0	119 ± 5	117 ± 11	116 ± 9
Biomass & ¹⁸ C Incorporation				
Total Biomass (Cells/bead)	5.68E+07	2.45E+05	1.92E+05	1.12E+05
¹³ C Enriched Biomass (Cells/bead)	4.98E+05	4.89E+03	2.47E+02	2.04E+01
Average PLFA Delta (‰)	255	245	198	432
Maximum PLFA Delta (‰)	1394	871	793	773
¹³ C Mineralization				
DIC Delta (‰)	38	-25	-13	-21
Community Structure (% total PLFA)				
Firmicutes (TerBrSats)	0.93	14.01	8.17	9.79
Proteobacteria (Monos)	36.67	53.17	69.86	57.59
Anaerobic metal reducers (BrMonos)	0.15	5.37	4.11	6.51
Actinomycetes (MidBrSats)	0.00	0.98	0.00	0.00
General (Nsats)	32.63	25.38	17.87	26.12
Eukaryotes (Polyenoics)	29.62	1.09	0.00	0.00
Physiological Status (Proteobacteria only)				
Slowed Growth	7.11	1.63	2.72	4.95
Decreased Permeability	0.03	0.14	0.02	0.00

Legend:

NA = Not analyzed NS = Not sampled J = Estimated result below PQL but above LQL I = Inhibited ND = Result not detected



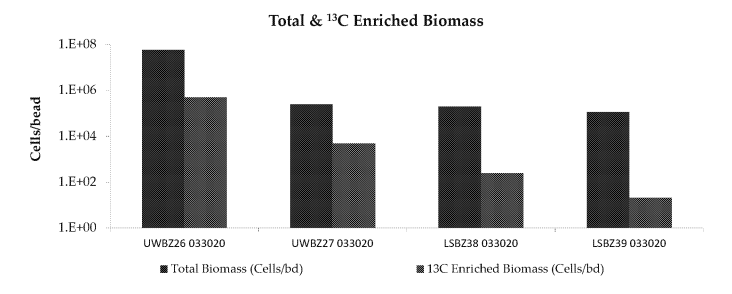


Figure 1. Biomass content is presented as a cell equivalent based on the total amount of phospholipid fatty acids (PLFA) extracted from a given sample. Total biomass is calculated based upon PLFA attributed to bacterial and eukaryotic biomass (associated with higher organisms).

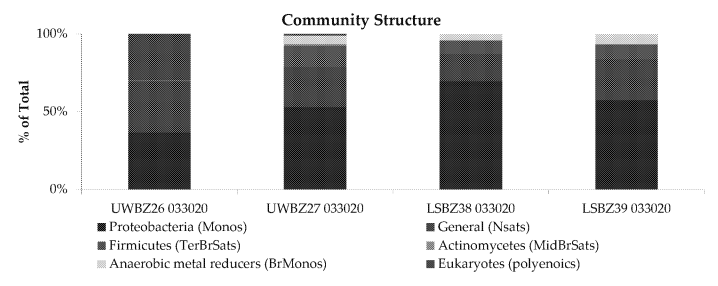


Figure 2. Relative percentages of total PLFA structural groups in the samples analyzed. Structural groups are assigned according to PLFA chemical structure, which is related to fatty acid biosynthesis. See the table in the interpretation section for detailed descriptions of the structural groups.



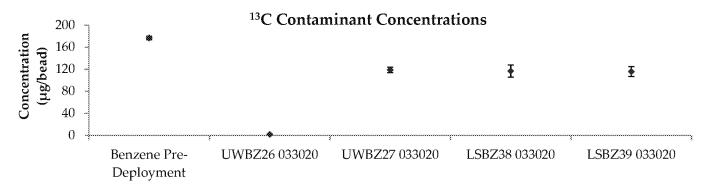


Figure 3. Comparison of Pre-deployment concentrations loaded on Bio-Sep beads to the concentrations detected after incubation.

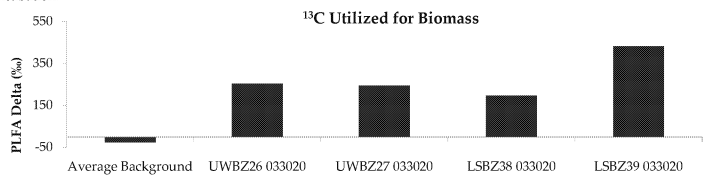


Figure 4. Comparison of the average Delta value obtained from PLFA biomarkers from each Bio-Trap® unit to the average background Delta observed in samples not exposed to ¹³C enriched compounds.

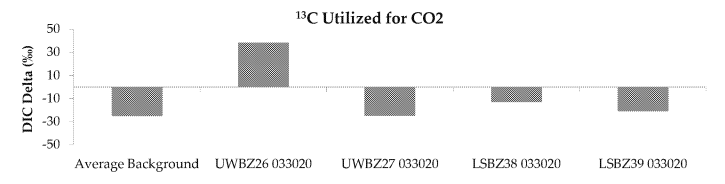


Figure 5. Comparison of the Delta value obtained from DIC from each Bio-Trap® unit to the average background Delta observed in samples not exposed to ¹³C enriched compounds.



Interpretation

Interpretation of the results of the SIP Bio-Trap® study must be performed with due consideration of site conditions, site activities, and the desired treatment mechanism. The following discussion describes interpretation of results in general terms and is meant to serve as a guide.

Contaminant Concentration: Bio-Traps® are baited with a ¹³C labeled contaminant of concern and a pre-deployment concentration is determined prior to shipping. Following deployment, Bio-Traps® are recovered for analysis including measurement of the concentration of the ¹³C labeled contaminant remaining. Pre- and post-deployment concentrations are used to calculate percent loss.

Biomass Concentrations: PLFA analysis is one of the most reliable and accurate methods available for the determination of viable (live) biomass. Phospholipids break down rapidly upon cell death (1,2), so biomass calculations based on PLFA content do not include "fossil" lipids from dead cells. Total biomass (cells/bead) is calculated from total PLFA using a conversion factor of 20,000 cells/pmole of PLFA. When making comparisons between wells, treatments, or over time, differences of one order of magnitude or more are considered significant.

Total Biomass			
Low	Moderate	High	
10³ to 10⁴ cells	10⁵ to 106 cells	10 ⁷ to 10 ⁸ cells	

¹³C Enriched Biomass: For SIP studies, ¹³C enriched PLFA is determined to quantify ¹³C incorporation into biomass as a line of evidence. The detection of ¹³C enriched biomass provides conclusive evidence of contaminant biodegradation. However, biodegradation of a contaminant of concern is almost always performed by a small subset of the total microbial community. Therefore, the ¹³C enriched biomass is typically several orders of magnitude lower than total biomass.

Average and Maximum PLFA Delta ¹³C: Isotopic data is often reported as a delta value. The delta value is the difference between the isotopic ratio (¹³C/¹²C) of the sample (R_x) and a standard (R_{std}) normalized to the isotopic ratio of the standard (R_{std}) and multiplied by 1,000 (units are parts per thousand or "per mill" and denoted ‰). R_{std} is the international standard Vienna PeeDee Belemnite (VPDB) with an anomalously high ¹³C/¹²C ratio of 0.011237. Due to the high value of the R_{std}, computed delta ¹³C values for most natural compounds are negative on a per mill basis.

Under natural conditions, the background delta 13 C value for PLFA is between -20 and -30‰. For a SIP Bio-Trap® study, biodegradation and incorporation of the 13 C labeled compound into PLFA results in a larger 13 C/ 12 C ratio (R_x) and thus delta values greater than under natural conditions.



Typical PLFA delta values are provided below.

PLFA Delta (‰)			
Low	Moderate	High	
0 to 100	100 to 1,000	>1,000	

Dissolved Inorganic Carbon (DIC): Often, bacteria can utilize the ¹³C labeled compound as both a carbon and energy source. The ¹³C portion used as a carbon source for growth can be incorporated into PLFA as discussed above, while the ¹³C used for energy is oxidized to ¹³CO₂ (mineralized).

 13 C enriched CO₂ data is often reported as a delta value as described above for PLFA. Under natural conditions, the delta 13 C value for CO₂ is typically in the range of -25% to -10% (3). For an SIP Bio-Trap® study, mineralization of the 13 C labeled contaminant of concern (increased 13 CO₂ production) would lead to a greater value of R_x and thus a positive delta value.

The detection of even low levels of ¹³C enriched DIC provides conclusive evidence of contaminant biodegradation. However, delta values between 0 and 100‰ are generally considered relatively low, values between 100 and 1,000‰ are considered moderate, and values greater than 1,000‰ are considered high.

Dissolved Inorganic Carbon (DIC) Delta and %13C			
Low	Moderate High		
0 to 100	100 to 1,000	>1,000	

Community Structure (% total PLFA): Community structure data is presented as a percentage of PLFA structural groups normalized to the total PLFA biomass. The relative proportions of the PLFA structural groups provide a "fingerprint" of the types of microbial groups (e.g. anaerobes, sulfate reducers, etc.) present and therefore offer insight into the dominant metabolic processes occurring at the sample location. Thorough interpretation of the PLFA structural groups depends in part on an understanding of site conditions and the desired microbial biodegradation pathways. For example, an increase in mid chain branched saturated PLFA (MidBrSats), indicative of sulfate reducing bacteria (SRB) and Actinomycetes, may be desirable at a site where anaerobic BTEX biodegradation is the treatment mechanism, but would not be desirable for a corrective action promoting aerobic BTEX or MTBE biodegradation. The following table provides a brief summary of each PLFA structural group and its potential relevance to bioremediation.



Description of PLFA structural groups.

Description of 1 2171 Structural groups.			
PLFA Structural Group	General classification	Potential Relevance to Bioremediation Studies	
Monoenoic (Monos)	Abundant in Proteobacteria (Gram negative bacteria), typically fast growing, utilize many carbon sources, and adapt quickly to a variety of	Proteobacteria is one of the largest groups of bacteria and represents a wide variety of both aerobes and anaerobes. The majority of Hydrocarbon utilizing bacteria fall within the	
1	environments.	Proteobacteria	
Terminally Branched Saturated (TerBrSats)	Characteristic of Firmicutes (Low G+C Grampositive bacteria), and also found in Bacteriodes, and some Gram-negative bacteria (especially anaerobes).	Firmicutes are indicative of presence of anaerobic fermenting bacteria (mainly <i>Clostridia/Bacteriodes</i> -like), which produce the H ₂ necessary for reductive dechlorination	
Branched Monoenoic (BrMonos)	Found in the cell membranes of micro-aerophiles and anaerobes, such as sulfate- or iron-reducing bacteria	In contaminated environments high proportions are often associated with anaerobic sulfate and iron reducing bacteria	
Mid-Chain Branched Saturated (MidBrSats)	Common in sulfate reducing bacteria and also Actinobacteria (High G+C Gram-positive bacteria).	In contaminated environments high proportions are often associated with anaerobic sulfate and iron reducing bacteria	
Normal Saturated (Nsats)	Found in all organisms.	High proportions often indicate less diverse populations.	
Polyenoic	Found in higher plants, and animals.	Eukaryotic scavengers will often prey on contaminant utilizing bacteria.	

Physiological Status (Proteobacteria): Some Proteobacteria modify specific PLFA as a strategy to adapt to stressful environmental conditions (4, 5). For example, *cis* monounsaturated fatty acids may be modified to cyclopropyl fatty acids during periods of slowed growth or modified to *trans* monounsaturated fatty acids to decrease membrane permeability in response to environmental stress. The ratio of product to substrate fatty acid thus provides an index of their health and metabolic activity.



Glossary

Delta (8): A Delta value is the difference between the isotopic ratio (13 C/ 12 C) of the sample (Rx) and a standard (Rstd) normalized to the isotopic ratio of the standard (Rstd) and multiplied by 1,000 (units are parts per thousand denoted %).

Delta = $(R_x-R_{std})/R_{std} \times 1000$

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